### Rev. January 2005

### **PATIENT INFORMATION LEAFLET Questions and Answers About Taxotere® Injection Concentrate**

(generic name = docetaxel) (pronounced as TAX-O-TEER)

### What is Taxotere?

Taxotere is a medication to treat breast cancer, nonsmall cell lung cancer and prostate cancer. It has severe side effects in some patients. This leaflet is designed to help you understand how to use Taxotere and avoid its side effects to the fullest extent possible. The more you understand your treatment, the better you will be able to participate in your care. If you have questions or concerns, be sure to ask your doctor or nurse. They are always your best source of information about your condition and treatment.

### What is the most important information about Taxotere?

- Since this drug, like many other cancer drugs, affects your blood cells, your doctor will ask for routine blood tests. These will include regular checks of your white blood cell counts. People with low blood counts can develop life-threatening infections. The earliest sign of infection may be fever, so if you experience a fever, tell your doctor right away.
- Occasionally, serious allergic reactions have occurred with this medicine. If you have any allergies, tell your doctor before receiving this medicine.
- A small number of people who take Taxotere have severe fluid retention, which can be life-threatening. To help avoid this problem, you must take another medication such as dexamethasone (DECKS-A-METH-A-SONE) prior to each Taxotere treatment. You must follow the schedule and take the exact dose of dexamethasone prescribed (see schedule at end of brochure). If you forget to take a dose or do not take it on schedule you must tell the doctor or nurse prior to your Taxotere treatment.
- If you are using any other medicines, tell your doctor before receiving your infusions of Taxotere.

## **How does Taxotere work?**

Taxotere works by attacking cancer cells in your body. Different cancer medications attack cancer cells in different ways.

Here's how Taxotere works: Every cell in your body contains a supporting structure (like a skeleton). Damage to this "skeleton" can stop cell growth or reproduction. Taxotere makes the "skeleton" in some cancer cells very stiff, so that the cells can no longer grow.

## **How will I receive Taxotere?**

Taxotere is given by an infusion directly into your vein. Your treatment will take about 1 hour. Generally, people receive Taxotere every 3 weeks. The amount of Taxotere and the frequency of your infusions will be determined by your doctor.

As part of your treatment, to reduce side effects your doctor will prescribe another medicine called dexamethasone. Your doctor will tell you how and when to take this medicine. It is important that you take the dexamethasone on the schedule set by your doctor. If you forget to take your medication, or do not take it on schedule, make sure to tell your doctor or nurse **BEFORE** you receive your Taxotere treatment. **Included with** this information leaflet is a chart to help you remember when to take your dexamethasone.

### What should be avoided while receiving Taxotere?

Taxotere can interact with other medicines. Use only medicines that are prescribed for you by your doctor and **be sure** to tell your doctor all the medicines that you use, including nonprescription drugs.

## What are the possible side effects of

**Low Blood Cell Count** – Many cancer medications, including Taxotere, cause a temporary drop in the number of white blood cells. These cells help protect your body from infection. Your doctor will routinely check your blood count and tell you if it is too low. Although most people receiving Taxotere do not have an infection even if they have a low white blood cell count, the risk of infection is increased.

Fever is often one of the most common and earliest signs of infection. Your doctor will recommend that you take your temperature frequently, especially during the days after treatment with Taxotere. If you have a fever, tell your doctor or nurse immediately.

**Allergic Reactions** – This type of reaction, which occurs during the infusion of Taxotere, is infrequent. If you feel a warm sensation, a tightness in your chest, or itching during or shortly after your treatment, tell your doctor or nurse immediately.

**Fluid Retention** – This means that your body is holding extra water. If this fluid retention is in the chest or around the heart it can be life-threatening. If you notice swelling in the feet and legs or a slight weight gain, this may be the first warning sign. Fluid retention usually does not start immediately; but, if it occurs, it may start around your 5th treatment. Generally, fluid retention will go away within weeks or months after your treatments are completed.

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**AXOTERE®** docetaxel **Injection Concentrate** 

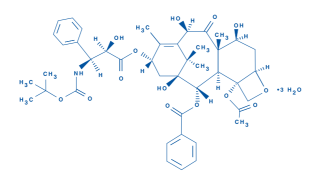
Prescribing Information as of January 2005 Rx only TAXOTERE®

TION LEA to Patient

FAXOTERE® (docetaxel) Injection Concentrate should be admin qualified physician experienced in the use of antineoplastic agents. Appropriate manag mplications is possible only when adequate diagnostic and treatment facilities are readily availab tompinations by possible only with adequate diagnostic and readment actuallies are treating available. The incidence of treatment-related mortality associated with TAXOTERE therapy is increased in patients with abnormal liver function, in patients receiving higher doses, and in patients with non-small cell lung carcinoma and a history of prior treatment with platinum-based chemotherapy who receive TAXOTERE as a single agent at a dose of 100 mg/m<sup>2</sup> (see **WARNINGS**). FAXOTERE should generally not be given to patients with bilirubin > upper limit of normal (ULN) for to patients with SGOT and/or SGPT >1.5 x ULN concomitant with alkaline phosphatase > 2.5 of to patients with soft analysis series and soft analysis of the prosphatase > 2.5 x ULN. Patients with elevations of bilirubin or abnormalities of transaminase concurrent with alkaline phosphatase are at increased risk for the development of grade 4 neutropenia, febrile neutropenia, infections, severe thrombocytopenia, severe stomatitis, severe skin toxicity, and toxic death. Patients with isolated elevations of transaminase > 1.5 x ULN also had a higher rate of febrile neutropenia grade 4 but did not have an increased incidence of toxic death. Bilirubin, SGOT or SGPT, and alkaline phosphatase values should be obtained prior to each cycle of AXOTERE therapy and reviewed by the treating physician.

AXOTERE therapy should not be given to patients with neutrophil counts of < 1500 cells/mm³. In order to monitor the occurrence of neutropenia, which may be severe and result in infection requent blood cell counts should be performed on all patients receiving TAXOTERE. vere hypersensitivity reactions characterized by hypotension and/or bronchospasm, or ger zed rash/erythema occurred in 2.2% (2/92) of patients who received the recommended 3-day dexa thasone premedication. Hypersensitivity reactions requiring discontinuation of the TAXOTER infusions preimentation, represensativity reactions requiring discontinuation of the TAX infusion were reported in five patients who did not receive premedication. These reactions re after discontinuation of the infusion and the administration of appropriate therapy. TAX must not be given to patients who have a history of severe hypersensitivity reactions to TAX or to other drugs formulated with polysorbate 80 (see WARNINGS). Severe fluid retention occurred in 6.5% (6/92) of patients despite use of a 3-day dexamethas remedication regimen. It was characterized by one or more of the following events: poorly toler-ted peripheral edema, generalized edema, pleural effusion requiring urgent drainage, dyspnea at sst, cardiac tamponade, or pronounced abdominal distention (due to ascites) (see **PRECAUTIONS**)

etaxel is an antineoplastic agent belonging to the taxoid family. It is prepared by semis



Docetaxel is a white to almost-white powder with an empirical formula of  $C_{43}H_{53}NO_{14}$ \* 3H<sub>2</sub>O, and a molecular weight of 861.9. It is highly lipophilic and practically insoluble in water. TAXOTERE (docetaxel) Injection Concentrate is a clear yellow to brownish-yellow viscous solution. TAXOTERE is sterile, non-pyrogenic, and is available in single-dose vials containing 20 mg (0.5 ml) or 80 mg (2 ml) docetaxel (anhydrous). Each mL contains 40 mg docetaxel (anhydrous) and 1040 mg polysorbate 80. trate requires dilution prior to use. A sterile, non-py

Docetaxel is an antineoplastic agent that acts by disrupting the microtubular network in cells that s essential for mitotic and interphase cellular functions. Docetaxel binds to free tubulin and promotes the assembly of tubulin into stable microtubules while simultaneously inhibiting their fisassembly. This leads to the production of microtubule bundles without normal function and to the stabilization of microtubules, which results in the inhibition of mitosis in cells. Docetaxel's ding to microtubules does not alter the number of protofilaments in the bound microtubules, a feature which differs from most spindle poisons currently in clinical use

The pharmacokinetics of docetaxel have been evaluated in cancer patients after administration of 20-115 mg/m² in phase I studies. The area under the curve (AUC) was dose proportional following doses of 70-115 mg/m² with infusion times of 1 to 2 hours. Docetaxel's pharmacokinetic profile is consistent with a three-compartment pharmacokinetic model, with half-lives for the  $\alpha$ ,  $\beta$ , and  $\gamma$ Consistent with a three-compariment pharmacokinetic mode, with nati-nives for the  $O_c$   $S_c$  and  $\gamma$  phases of 4 min, 36 min, and 11.1 hr, respectively. The initial rapid decline represents distribution to the peripheral compartments and the late (terminal) phase is due, in part, to a relatively slow efflux of docetaxel from the peripheral compartment. Mean values for total body clearance and steady state volume of distribution were 21  $L/h/m^2$  and 113 L, respectively. Mean total body clearance for Japanese patients dosed at the range of 10-90 mg/m² was similar to that of European/American populations dosed at 100 mg/m², suggesting no significant difference in the elimination of docetaxel in the two populations.

is study of <sup>14C</sup>-docetaxel was conducted in three cancer patients. Docetaxel was eliminated in both the urine and feces following oxidative metabolism of the *tert-*butvl exter group, but facel programs. axer was conducted in three carder patients. Doctaxer was eminiated in our following oxidative metabolism of the tert-butyl ester group, but fecal excretion nation route. Within 7 days, urinary and fecal excretion accounted for approximation route. vered in feces is excreted during the first 48 hours as 1 major and 3 minor metabolites with ver small amounts (less than 8%) of unchanged drug.

A population pharmacokinetic analysis was carried out after TAXOTERE treatment of 535 patients dosed at 100 mg/m². Pharmacokinetic parameters estimated by this analysis were very close to those estimated from phase I studies. The pharmacokinetics of docetaxel were not influenced by age or gender and docetaxel total body clearance was not modified by pretreatment with dexamet and/or SePT > 1.5 times the upper filmt of normal JULNJ contominant with alialine phosphala > 2.5 times ULNJ, total body clearance was lowered by an average of 27%, resulting in a 38% increase systemic exposure (AUC). This average, however, includes a substantial range and there is, at preser no measurement that would allow recommendation for dose adjustment in such patients. Patien rmalities of transaminase and alkaline phosphatase should, in general, not b

Clearance of docetaxel in combination therapy with cisplatin was similar to that previously observe Clearance of docetaxel in combination therapy with cisplatin was similar to that previously observed following monotherapy with docetaxel. The pharmacokinetic profile of cisplatin in combination therapy with docetaxel was similar to that observed with cisplatin alone.

A population pharmacokinetic analysis of plasma data from 40 patients with hormone-refractory metastatic prostate cancer indicated that docetaxel systemic clearance in combination with prednisone is similar to that observed following administration of docetaxel alone. A study was conducted in 30 patients with advanced breast cancer to determine the potential for drug-drug-interactions between docetaxel (75 mg/m²), doxorubicin (50 mg/m²), and cyclophosphamide (500 mg/m²) when administered in combination. The coadministration of docetaxel had no effect on the pharmacokinetics of doxorubicin and cyclophosphamide when the three drugs vere given in combination compared to coadministration of doxorubicin and cyclopho only. In addition, doxorubicin and cyclophosphamide had no effect on docetaxel plasma clearance when the three drugs were given in combination compared to historical data for docetaxe

vitro studies showed that docetaxel is about 94% protein bound, mainly to  $lpha_1$ -acid glyco albumin, and lipoproteins. In three cancer patients, the *in vitro* binding to plasma proteins was found to be approximately 97%. Dexamethasone does not affect the protein binding of docetaxel. In vitro drug interaction studies revealed that docetaxel is metabolized by the CYP3A4 isoenzyme and its metabolism can be inhibited by CYP3A4 inhibitors, such as ketoconazole, erythromycin troleandomycin, and nifedipine. Based on *in vitro* findings, it is likely that CYP3A4 inhibitors and/or substrates may lead to substantial increases in docetaxel blood concentrations. No clinical studies have been performed to evaluate this finding (see **PRECAUTIONS**).

Breast Cancer
The efficacy and safety of TAXOTERE have been evaluated in locally advanced or metastatic breast cancer after failure of previous chemotherapy (alkylating agent-containing regimens or anthracy-**Randomized Trials** 

In one randomized trial, patients with a history of prior treatment with an anthracycline-contain men were assigned to treatment with TAXOTERE (100 mg/m² every 3 weeks) or the combination intomycin (12 mg/m² every 6 weeks) and vinblastine (6 mg/m² every 3 weeks). 203 patients were fomized to TAXOTERE and 189 to the comparator arm. Most patients had received prior motherapy for metastatic disease; only 27 patients on the TAXOTERE arm and 33 patients on the comparator arm. imparator arm entered the study following relapse after adjuvant therapy. Three-quarters of ents had measurable, visceral metastases. The primary endpoint was time to progression. The

## Efficacy of TAXOTERE in the Treatment of Breast Cancer Patients Previously Treated with an Anthracycline-Containing Regimen (Intent-to-Treat Analysis)

Efficacy Parameter	Docetaxel Mitomycin/ Vinblastine		p-value
	(n=203)	(n=189)	
Median Survival	11.4 months	8.7 months	
Risk Ratio*, Mortality (Docetaxel: Control)	0.	.73	p=0.01 Log Rank
95% CI (Risk Ratio)	0.58	3-0.93	
Median Time to Progression	4.3 months	2.5 months	
Risk Ratio*, Progression (Docetaxel: Control)	0.	p=0.01 Log Rank	
95% CI (Risk Ratio)	0.61	-0.94	
Overall Response Rate Complete Response Rate	28.1% 3.4%	9.5% 1.6%	p<0.0001 Chi Square

assigned to treatment with TAXOTERE (100 mg/m²) or doxorubicin (75 mg/m²) every 3 weeks. 16 patients were randomized to TAXOTERE and 165 patients to doxorubicin. Approximately one-half of nations had received prior chemotherapy for metastatic disease, and one-half entered the study ollowing relapse after adjuvant therapy. Three-quarters of patients had measurable, visceral metas-ases. The primary endpoint was time to progression. The study results are summarized below:

## **Efficacy of TAXOTERE in the Treatment of Breast Cancer Patients**

Previously Treated with an Alkylating-Containing Regimen (Intent-to-Treat Analysis)					
Efficacy Parameter	Docetaxel (n=161)	p-value			
Median Survival	14.7 months	14.3 months			
Risk Ratio*, Mortality (Docetaxel: Control)	0.	p=0.39 Log Rank			
95% CI (Risk Ratio)	0.68-1.16				
Median Time to Progression	6.5 months	5.3 months			
Risk Ratio*, Progression (Docetaxel: Control)	0.	p=0.45 Log Rank			
95% CI (Risk Ratio)	0.71	-1.16			
Overall Response Rate Complete Response Rate	45.3% 6.8%	29.7% 4.2%	p=0.004 Chi Square		

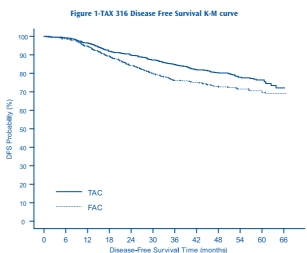
In another multicenter open-label, randomized trial (TAX313), in the treatment of patients with advanced breast cancer who progressed or relapsed after one prior chemotherapy regimen, 527 patients were randomized to receive TAXOTERE monotherapy 60 mg/m² (n=151), 75 mg/m² (n=188) or 100 mg/m<sup>2</sup> (n=188). In this trial, 94% of patients had metastatic disease and 79% had received prior anthracycline therapy. Response rate was the primary endpoint. Response rates increased with TAXOTERE dose: 19.9% for the 60 mg/m² group compared to 22.3% for the 75 mg/m² and 29.8% for the 100 mg/m² group; pair-wise comparison between the 60 mg/m² and 100 mg/m² groups was ally significant, (p=0.037).

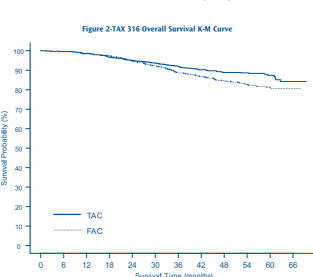
TAXOTERE at a dose of 100 mg/m<sup>2</sup> was studied in six single arm studies involving a total of 309 ients with metastatic breast cancer in whom previous chemotherapy had failed. Among these, 190 ients had anthracycline-resistant breast cancer, defined as progression during an anthracycline-taining chemotherapy regimen for metastatic disease, or relapse during an anthracyclinening adjuvant regimen. In anthracycline-resistant patients, the overall response rate was 37.9% (72/190; 95% C.I.: 31.0-44.8) and the complete response rate was 2.1%. AXOTERE was also studied in three single arm Japanese studies at a dose of 60 mg/m², in 174 NAOTERE was also studied in three single arm Japanese studies at a dose of 60 mg/m², in 174 Datients who had received prior chemotherapy for locally advanced or metastatic breast cancer. Among 26 patients whose best response to an anthracycline had been progression, the response rate was 34.6% (95% C.L.: 17.2-55.7), similar to the response rate in single arm studies of 100 mg/m².

**Adjuvant Treatment of Breast Cancer** nter, open-label, randomized trial (TAX316) evaluated the efficacy and safety of TAXOTERE for the adjuvant treatment of patients with axillary-node-positive breast cancer and no evidence followed by fluorouracil 500 mg/m<sup>2</sup> and cyclosphosphamide 500 mg/m<sup>2</sup> (FAC arm). Both regin nistered every 3 weeks for 6 cycles. TAXOTERE was admir were administered every 3 weeks for 6 tytues. ITANIERL was administered every 3 weeks for 6 tytues. ITANIERL was administered every 3 weeks for 6 tytues. ITANIERL was administered every 3 weeks for 6 tytues. ITANIERL was administered to 10 the modern and for progestering received tamoxifen 20 mg daily for patients with positive estrogen and/or progesterone receptors received tamoxifen 20 mg daily for up to 5 years. Adjuvant radiation therapy was prescribed according to guidelines in place at participating institutions and was given to 69% of patients who received TAC and 72% of patients who Results from a second interim analysis (median follow-up 55 months) are as follows: In study TAX

316, the docetaxel-containing combination regimen TAC showed significantly longer disease-free survival (DFS) than FAC (hazard ratio=0.74; 2-sided 95% CI=0.60, 0.92, stratified log rank p=0.0047). The primary endpoint, disease-free survival, included local and distant recurrences, contralateral breast cancer and deaths from any cause. The overall reduction in risk of relapse was 25.7% for TAC-At the time of this interim analysis, based on 219 deaths, overall survival was longer for TAC than FAC (hazard ratio=0.69, 2-sided 95% CI=0.53, 0.90). (See Figure 2). There will be further analysis at

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		Disease	Free Survival	0verall	Survival
Patient subset	Number of patients	Hazard ratio*	95% CI	Hazard ratio*	95% CI
No. of positive nodes					
Overall	744	0.74	(0.60, 0.92)	0.69	(0.53, 0.90)
1-3	467	0.64	(0.47, 0.87)	0.45	(0.29, 0.70)
4+	277	0.84	(0.63, 1.12)	0.93	(0.66, 1.32)
Receptor status					
Positive	566	0.76	(0.59, 0.98)	0.69	(0.48, 0.99)
Negative	178	0.68	(0.48, 0.97)	0.66	(0.44, 0.98)

Non-Small Cell Lung Cancer (NSCLC) The efficacy and safety of TAXOTERE has been evaluated in patients with unresectable, locall

advanced or metastatic non-small cell lung cancer whose disease has failed prior platinum-based Monotherapy with TAXOTERE for NSCLC Previously Treated with Platinum-Based Chemo Two randomized, controlled trials established that a TAXOTERE dose of 75 mg/m² was toler yielded a favorable outcome in patients previously treated with platinum-based chemothe below). TAXOTERE at a dose of 100 mg/m², however, was associated with unacceptable her ted mortality and this dose should not be used (see BOXED WARNING, WARNINGS, and DOSAGE AND ADMINISTRATION sections).

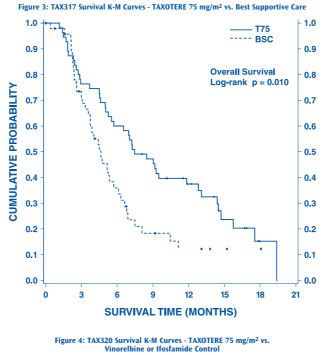
One trial (TAX317), randomized patients with locally advanced or metastatic non-small cell lung cancer, a history of prior platinum-based chemotherapy, no history of taxane exposure, and an ECOG performance status \$2\text{ to TAXOTERE or loest supportive care. The primary endpoint of the study was survival. Patients were initially randomized to TAXOTERE 100 mg/m² or best supportive care, but early toxic deaths at this dose led to a dose reduction to TAXOTERE 75 mg/m<sup>2</sup>. A total of 104 patients were randomized in this amended study to either TAXOTERE 75 mg/m2 or bes a second randomized trial (TAX320), 373 patients with locally advanced or metastatic non-sm

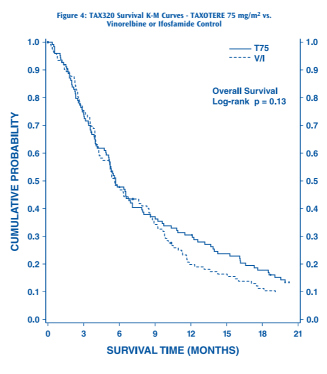
cell lung cancer, a history of prior platinum-based chemotherapy, and an ECOG performance status \$2 were randomized to TAXOTERE 75 mg/m², TAXOTERE 100 mg/m² and a treatment in which the investigator chose either vinorelbine 30 mg/m² days 1, 8, and 15 repeated every 3 weeks on the contract of the contr ifosfamide 2 g/m² days 1-3 repeated every 3 weeks. Forty percent of the patients in this study had history of prior paclitaxel exposure. The primary endpoint was survival in both trials. The efficat data for the TAXOTERE 75 mg/m² arm and the comparator arms are summarized in the table below and in figures 3 and 4 showing the survival curves for the two studies.

Efficacy of TAXOTERE in the Treatment of Non-Small Cell Lung Cancer Patients Previously Treated with a Platinum-Based Chemotherapy Regimen (Intent-to-Treat Analysis)
TAX317 TAX320

	Docetaxel 75 mg/m² n=55	Best Supportive Care/75 n=49	Docetaxel 75 mg/m² n=125	Control (V/I) n=123
Overall Survival Log-rank Test	p=0	).01	p=	0.13
Risk Ratio <sup>††</sup> , Mortality (Docetaxel: Control) 95% CI (Risk Ratio)	0.! (0.35,			82 , 1.06)
Median Survival 95% CI	7.5 months* (5.5, 12.8)	4.6 months (3.7, 6.1)	5.7 months (5.1, 7.1)	5.6 months (4.4, 7.9)
% 1-year Survival 95% CI	37%*† (24, 50)	12% (2, 23)	30%*† (22, 39)	20% (13, 27)
Time to Progression 95% CI	12.3 weeks* (9.0, 18.3)	7.0 weeks (6.0, 9.3)	8.3 weeks (7.0, 11.7)	7.6 weeks (6.7, 10.1)
Response Rate 95% CI	5.5% (1.1, 15.1)	Not Applicable	5.7% (2.3, 11.3)	0.8% (0.0, 4.5)

Only one of the two trials (TAX317) showed a clear effect on survival, the primary en trial also showed an increased rate of survival to one year. In the second study (TAX320) the rate of survival at one year favored TAXOTERE 75 mg/m<sup>2</sup>





Patients treated with TAXOTERE at a dose of 75 mg/m<sup>2</sup> experienced no deterioration in performance

status and body weight relative to the comparator arms used in these trials.

Combination Therapy with TAXOTERE for Chemotherapy-Naïve NSCLC

In a randomized controlled trial (TAX326), 1218 patients with unresectable stage IIIB or IV NSCLC and no prior chemotherapy were randomized to receive one of three treatments:

TAXOTERE 75 mg/m² as a 1 hour infusion immediately followed by cisplatin 75 mg/m² over 30-60 in the control of the control utes every 3 weeks; vinorelbine 25 mg/m<sup>2</sup> administered over 6-10 minutes on days 1, 8, 15, 22 followed by cisplatin 100 mg/m<sup>2</sup> administered on day 1 of cycles repeated every 4 weeks; or a

The primary efficacy endpoint was overall survival. Treatment with TAXOTERE+cisplatin did not result in a statistically significantly superior survival compared to vinorelbine+cisplatin (see table below). The 95% confidence interval of the heazard ratio (adjusted for interim analysis and multiple risons) shows that the addition of TAXOTERE to cisplatin results in an outcome ranging fro a 6% inferior to a 26% superior survival compared to the addition of vinorelbine to cisp results of a further statistical analysis showed that at least (the lower bound of the 95% confider

Comparison	Taxotere+Cisplatin n=408	Vinorelbine+Cisplatin n=405
Kaplan-Meier Estimate of Median Survival	10.9 months	10.0 months
p-value <sup>a</sup>	0.	122
Estimated Hazard Ratiob	0	.88
Adjusted 95% CI <sup>c</sup>	(0.74	, 1.06)

that TAXOTERE+cisplatin is associated with a longer survival The second comparison in the study vinorelline+cisplatin versus TAXOTERE+carbonlatin, did not

the second companisor in the study, kindroline-topicalin versus TAXOTERE-Carobipathi, but not demonstrate superior survival associated with the TAXOTERE arm (Kaplan-Meier estimate of median survival was 9.1 months for TAXOTERE+carboplatin compared to 10.0 months on the vinorelbine+cisplatin arm) and the TAXOTERE+carboplatin arm did not demonstrate preservation of at least 50% of the survival effect of vinorelbine added to cisplatin. Secondary endpoints evalu ignificant difference between TAXOTERE+cisplatin and vive response and time to progression (see table below).

# Response and TTP Analysis of TAXOTERE in Combination Therapy for

Endpoint	TAXOTERE+ Cisplatin	Vinorelbine+ Cisplatin	p-value
Objective Response Rate	31.6%	24.4%	Not Significant
(95% CI)a	(26.5%, 36.8%)	(19.8%, 29.2%)	Significant
Median Time to Progression <sup>b</sup>	21.4 weeks	22.1 weeks	Not Significant
(95% CI)a	(19.3, 24.6)	(18.1, 25.6)	

bKaplan-Meier estimate

endent (hormone refractory) metastatic prostate cancer were evaluated in a rando trol trial. A total of 1006 patients with Karnofsky Performance Status (KPS) ≥6

were randomized to the following treatment groups:

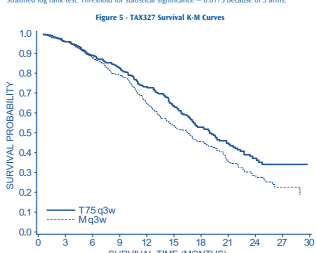
TAXOTERE 75 mg/m² every 3 weeks for 10 cycles.

TAXOTERE 30 mg/m² administered weekly for the first 5 weeks in a 6-week cycle for 5 cycles.

Mitoxantrone 12 mg/m² every 3 weeks for 10 cycles. All 3 regimens were administered in combination with prednisone 5 mg twice daily, continu n the TAXOTERE every three week arm, a statistically significant overall survival advantage wa ne. In the TAXOTERE weekly arm, no overall survival a

Efficacy of TAXOTERE in the Treatment of Patients with Androgen Independen

	every 3 weeks	Mitoxantrone every 3 weeks
Number of patients	335	337
Median survival (months)	18.9	16.5
95% CI	(17.0-21.2)	(14.4-18.6)
Hazard ratio	0.761	
95% CI	(0.619-0.936)	
p-value*	0.0094	



INDICATIONS AND USAG

TAXOTERE is indicated for the treatment of patients with locally advanced or metastatic breast TAXOTERE in combination with doxorubicin and cyclophosphamide is indicated for the adjuvant treatment of patients with operable node-positive breast cancer.

Non-Small Cell Lung Cancer

TAXOTER: each of the Adjuvant for the Adj

TAXOTERE in combination with cisplatin is indicated for the treatment of patients with u locally advanced or metastatic non-small cell lung cancer who have not previously received

TAXOTERE in combination with prednisone is indicated for the treatment of patients with androgen independent (hormone refractory) metastatic prostate cancer

## CONTRAINDICATIONS

ndicated in patients who have a history of severe hypersensitivity reactions to

ate diagnostic and treatment facilities are readily available

TAXOTERE administered at 100 mg/m² was associated with deaths considered possibly or probably related to treatment in 2.0% (19/965) of metastatic breast cancer patients, both previously treated and untreated, with normal baseline liver function and in 11.5% (7/61) of patients with various nor types who had abnormal baseline liver function (SGOT and/or SGPT > 1.5 times ULN togethe with AP > 2.5 times ULN). Among patients dosed at 60 mg/m², mortality related to treatme occurred in 0.6% (3/481) of patients with normal liver function, and in 3 of 7 patients with abnorm Non-Small Cell Lung Cancer

small cell lung cancer who had a history of prior platinum-based chemotherapy was associated with infance in ling cancer with had a instory of prior plantifull-based chemometapy was associated with creased treatment-related mortality (14% and 5% in two randomized, controlled studies). There ere 2.8% treatment-related deaths among the 176 patients treated at the 75 mg/m² dose in the indomized trials. Among patients who experienced treatment-related mortality at the 75 mg/m² dose level, 3 of 5 patients had a PS of 2 at study entry (see **BOXED WARNING, CLINICAL STUDIES**, and DOSAGE AND ADMINISTRATION sections).

uid be premedicated with oral corticosterous (see below for prostate cancer) such 16 mg per day (e.g., 8 mg BID) for 3 days starting 1 day prior to TAXOTERE to redi fluid retention and hypersensitivity reactions (see **DOSAGE AND ADMINISTRATI**) section). This regimen was evaluated in 92 patients with metastatic breast cancer previously treater with chemotherapy given TAXOTERE at a dose of 100 mg/m<sup>2</sup> every 3 weeks.

e 8 mg, at 12 hours, 3 hours and 1 hour before the TAXOTERE infusion (see DOSAGE AND Patients should be observed closely for hypersensitivity reactions, especially during the first an

second infusions. Severe hypersensitivity reactions characterized by hypotension and/o bronchospasm, or generalized rash/erythema occurred in 2.2% of the 92 patients premedicated with vity reactions should not be rechallenged with TAXOTERE.

< 2000 neutrophils/mm³) occurs in virtually all patients given 60-100 mg/m² of</p> TAXOTERE and grade 4 neutropenia (< 500 cells/mm³) occurs in 85% of patients given 100 mg/m and 75% of patients given 60 mg/m². Frequent monitoring of blood counts is, therefore, essential st that dose can be adjusted. TAXOTERE should not be administered to patients with neutrophils <

Febrile neutropenia occurred in about 12% of patients given 100 mg/m<sup>2</sup> but was very und nents given 60 mg/m². Hematologic responses, febrile reactions and infections, and rates of septic th for different regimens are dose related and are described in **CLINICAL STUDIES**. se breast cancer patients with severe liver impairment (bilirubin > 1.7 times ULN) developed

cute Myeloid Leukemia reatment-related acute myeloid leukemia (AML) has occurred in patients given anthracyclines moldor cyclophosphamide, including use in adjuvant therapy for breast cancer. In the adjuvant reast cancer trial (TAX316, see CLINICAL STUDIES) AML occurred in 3 of 744 patients who received TAXOTERE, doxorubicin and cyclophosphamide and in 1 of 736 patients who received fluorouraci orubicin and cyclophosphamide (see ADVERSE REACTIONS).

rabbits at doses ≥ 0.3 and 0.03 mg/kg/day, respectively (about 1/50 and 1/300 the daily maximu recommended human dose on a mg/m<sup>2</sup> basis), administered during the period of organogenesis have shown that TAXOTERE is embryotoxic and fetotoxic (characterized by intrauterine mortality creased resorption, reduced fetal weight, and fetal ossification delay). The doses indicated above so caused maternal toxicity.

ere are no adequate and well-controlled studies in pregnant women using TAXOTERE. If TAXOTERE is used during pregnancy, or if the patient becomes pregnant while receiving this drug the patient should be apprised of the potential hazard to the fetus or potential risk for loss of the pregnancy. Women of childbearing potential should be advised to avoid becoming pregnant during

Responding patients may not experience an improvement in performance status on therapy and may experience worsening. The relationship between changes in performance status, response to therapy, and treatment-related side effects has not been established.

In order to monitor the occurrence of myelotoxicity, it is recommended that frequency perspectable blood cell counts be performed on all patients receiving TAXOTERE. Patients should not be retreated with subsequent cycles of TAXOTERE until neutrophils recover to a level > 1500 cells/mm³ and platelets recover to a level > 100,000 cells/mm³.

troin in a TAXOTERE cycle (see **DOSAGE AND ADMINISTRATION** section).

Hypersensitivity **Reactions**Hypersensitivity reactions may occur within a few minutes following initiation of a TAXOTERE infusion. If minor reactions such as flushing or localized skin reactions occur, interruption of therapy is not required. More severe reactions, however, require the immediate discontinuation of OTERE and aggressive therapy. All patients should be premedicated with an oral corti ion of the infusion of TAXOTERE (see BOXED WARNING and WARNINGS: Premed-

Localized erythema of the extremities with edema followed by desquamation has be case of severe skin toxicity, an adjustment in dosage is recommended (see **DOSAGE AND ADMINIS-TRATION** section). The discontinuation rate due to skin toxicity was 1.6% (15/965) for metastatic ast cancer patients. Among 92 breast cancer patients premedicated with 3-day corticosteroi we were no cases of severe skin toxicity reported and no patient discontinued TAXOTERE due

ention has been reported following TAXOTERE therapy (see BOXED WARNING and WARNINGS: Premedication Regimen). Patients should be premedicated with oral cortic WARNINGS: Premedication Regimen). Fatents should be premedicated with oral corticosteroids prior to each TAXOTERE administration to reduce the incidence and severity of fluid retention (see DOSAGE AND ADMINISTRATION section). Patients with pre-existing effusions should be closely monitored from the first dose for the possible exacerbation of the effusions. When fluid retention occurs, peripheral edema usually starts in the lower extremities and may become generalized with a median weight gain of 2 kg.

occurred in 27.2% and severe fluid retention in 6.5%. The median cumulative dose to onset of moderate or severe fluid retention was 819 mg/m². 9.8% (9/92) of patients discontinued treatment due to fluid retention: 4 patients discontinued with severe fluid retention; 4 patients discontinued with severe fluid retention; 4 patients discontinued with severe fluid retention; the remaining 5 had mild or moderate fluid retention. The median cumulative dose to treatment discontinuation due to fluid redian of 16 weeks from the last infusion of TAXOTERE to resolution (range: 0 to 42+ weeks)

sory symptoms (paresthesia, dysesthesia, pain) were observed in 5.5% (53/965) of etastatic breast cancer patients, and resulted in treatment discontinuation in 6.1%. When thes ir, dosage must be adjusted. If symptoms persist, treatment should be disco promo occur, uoage must be adjusted. In symptoms persist, treatment should be discontinued e DOSAGE AND ADMINISTRATION section). Patients who experienced neurotoxicity in clinical als and for whom follow-up information on the complete resolution of the event was available d spontaneous reversal of symptoms with a median of 9 weeks from onset (range; 0 to 106 weeks). Severe peripheral motor neuropathy mainly manifested as distal extremity weakness occurred in

Severe asthenia has been reported in 14.9% (144/965) of metastatic breast cancer patients but has led to treatment discontinuation in only 1.8%. Symptoms of fatigue and weakness may last a few days up to several weeks and may be associated with deterioration of performance status in patients with progressive disease.

Information for Patients

ation, see the accompanying Patient Information Leaflet. Drug Interactions
There have been no formal clinical studies to evaluate the drug interactions of TAXOTERE with other medications. *In vitro* studies have shown that the metabolism of docteaxel may be modified by the medications.

itant administration of compounds that induce, inhibit, or are metabolized by cytochron P450 3A4, such as cyclosporine, terfenadine, ketoconazole, erythromycin, and troles ion should be exercised with these drugs when treating patients receiving TAXOTERE as there i Carcinogenicity, Mutagenicity, Impairment of Fertility

No studies have been conducted to assess the carcinogenic potential of TAXOTERE. TAXOTERE has been shown to be clastogenic in the *in vitro* chromosome aberration test in CHO-K<sub>1</sub> cells and in the

in vivo micronucleus test in the mouse, but it did not induce mutagenicity in the Ames test or the CHO/HGPRT gene mutation assays TAXOTERE produced no impairment of fertility in rats who

administered in multiple IV doses of up to 0.3 mg/kg (about 1/50 the recommended human dose on a mg/m² basis), but decreased testicular weights were reported. This correlates with findings of a 10-cycle toxicity study (dosing once every 21 days for 6 months) in rats and dogs in which testicular atrophy or degeneration was observed at IV doses of 5 mg/kg in rats and 0.375 mg/kg in dogs (about 1/3 and 1/15 the recommended human dose on a mg/m² basis, respectively). An increference of dosing in rats produced similar effects at lower dose levels.

Pregnancy
Pregnancy Category D (see WARNINGS section). It is not known whether TAXOTERE is excreted in human milk. Because many drugs are excreted in human milk, and because of the potential for serious adverse reactions in nursing infants from TAXOTERE, mothers should discontinue nursing prior to taking the drug.

d effectiveness of TAXOTERE in pediatric patients have not been established

In a study conducted in chemotherapy-naïve patients with NSCLC (TAX326), 148 patients (36%) in the norelbine+cisplatin group 65 years of age or greater. In the TAXOTERE+cisplatin group, pss than 65 years of age had a median survival of 10.3 months (95% CI : 9.1 months, 11.8 m dp atients 65 years or older had a median survival of 12.1 months (95% CI : 9.3 month onths). In patients 65 years or fage or greater treated with TAXOTERE+cisplatin, diarrhea peripheral edema (39%) and stomatitis (28%) were observed more frequently than in the vinc bine+cisplatin group (diarrhea 24%, peripheral edema 20%, stomatitis 20%). Patients treated with TAXOTERE+cisplatin who were 65 years of age or greater were more likely to experience diarrhea (55%), infections (42%), peripheral edema (39%) and stomatitis (28%) compared to patients less than ID number: 50074275 Version Country USA Date: 2-2-05 Operator: Logo version: SCV A2 Product Desc. Insert, Taxotere, 80mg/20mg Dagenham Supplier: Artwork created in: QuarkXpress 4.0, Mac format Fonts used: OceanSansAV Light, Bold, Light Italic Minimum point size of text: 5 pt. Colors Used: Reflex Blue

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When TAXOTERE was combined with carboplatin for the treatment of chemotherapy-naïve oma, patients 65 years of age or greater (28%) experienced by the similar patients treated with TAXOTERE+cisplatin, and a nced non-small cell lung carcino TVANCED RODA-SHAIL CELLUING CALCIDINA, PARIETIS DO YEARS OF AGE OF BECAUTE (2009) Experiences gipher frequency of infection compared to similar patients treated with TAXOTERE+cisplatin, and a igher frequency of diarrhea, infection and peripheral edema than elderly patients treated with Vinoreibine+cispiatin.

Of the 333 patients treated with TAXOTERE every three weeks plus prednisone in the prostate ca tudy (TAX327), 209 patients were 65 years of age or greater and 68 patients were older than 75 ears. In patients treated with TAXOTERE every three weeks, the following TEAEs occurred at rates ≥ 10% higher in patients 65 years of age or greater compared to younger patients: anemia (71% vs 59%), infection (37% vs. 24%), nail changes (34% vs. 23%), anorexia (21% vs. 10%), weight loss (15% vs n the adjuvant breast cancer trial (TAX316), TAXOTERE in combination with doxorubicin and cyclophosphamide was administered to 744 patients of whom 48 (6%) were 65 years of age or greater. The number of elderly patients who received this regimen was not sufficient to determine whether there were differences in safety and efficacy between elderly and younger patients.

erse reactions are described for TAXOTERE according to indication

in the treatment of breast cancer, at the maximum dose of 100 mg/m<sup>2</sup> in the treatment of advanced breast cancer at doses of 60, 75 and 100 mg/m<sup>2</sup> in the adjuvant therapy of breast cancer at dose of 75 mg/m<sup>2</sup>, in combination with doxorubicin ment of advanced non-small cell lung cancer after prior platinum-based chemotherapy, nent of non-small cell lung cancer in patients who have not previously received therapy for this condition, at a dose of 75 mg/m<sup>2</sup>, in combination with cisplatin treatment of androgen independent (hormone refractory) metastatic prostate cancer, at a rany with TAXOTERE for Locally Advanced or Metastatic Breast Cancer After Failure

of Prior Chemotherapy
TAXOTERE 100 mg/m<sup>2</sup>: Adverse drug reactions occurring in at least 5% of patients are compared for three populations who received TAXOTERE administered at 100 mg/m² as a 1-hour infusion every 3 weeks: 2045 patients with various tumor types and normal baseline liver function tests; the subset of 965 patients with locally advanced or metastatic breast cancer, both previously treated and untreated with chemotherapy, who had normal baseline liver function tests; and an additional 61 atients with various tumor types who had abnormal liver function tests at base ere described using COSTART terms and were considered possibly or probably related to TAXOTER! nilar in patients receiving TAXOTERE for the treatment of breast cancer and in patients with other

Summary of Adverse Events in Patients Receiving TAXOTERE at 100 mg/m<sup>2</sup>

Summary of Autresse	All Tumor	All Tumor	Breast
Adverse Event	Types Normal LFTs* n=2045	Types Elevated LFTs** n=61 %	Cancer Normal LFTs* n=965
Hematologic	/0	/0	/0
Neutropenia			
<2000 cells/mm <sup>3</sup>	95.5	96.4	98.5
<500 cells/mm <sup>3</sup>	75.4	87.5	85.9
Leukopenia	05.6	00.2	00.6
<4000 cells/mm <sup>3</sup> <1000 cells/mm <sup>3</sup>	95.6 31.6	98.3 46.6	98.6 43.7
Thrombocytopenia	31.0	40.0	45.7
<100,000 cells/mm <sup>3</sup>	8.0	24.6	9.2
Anemia			
<11 g/dL	90.4	91.8	93.6
<8 g/dL	8.8	31.1	7.7
Febrile Neutropenia***	11.0	26.2	12.3
Septic Death Non-Septic Death	1.6 0.6	4.9 6.6	1.4 0.6
Infections	0.0	0.0	0.0
Any	21.6	32.8	22.2
Severe	6.1	16.4	6.4
Fever in Absence of Infection			
Any	31.2	41.0	35.1
Severe	2.1	8.2	2.2
Hypersensitivity Reactions			
Regardless of Premedication Any	21.0	19.7	17.6
Severe	4.2	9.8	2.6
With 3-day Premedication	n=92	n=3	n=92
Any	15.2	33.3	15.2
Severe	2.2	0	2.2
Fluid Retention			
Regardless of Premedication	47.0	39.3	59.7
Any Severe	6.9	8.2	8.9
With 3-day Premedication	n=92	n=3	n=92
Any	64.1	66.7	64.1
Severe	6.5	33.3	6.5
Neurosensory			
Any	49.3	34.4	58.3
Severe Cutaneous	4.3	0	5.5
Any	47.6	54.1	47.0
Severe	4.8	9.8	5.2
Nail Changes			
Any	30.6	23.0	40.5
Severe	2.5	4.9	3.7
Gastrointestinal			
Nausea	38.8 22.3	37.7 23.0	42.1 23.4
Vomiting Diarrhea	22.3 38.7	32.8	42.6
Severe	4.7	4.9	5.5
Stomatitis			
Any	41.7	49.2	51.7
Severe	5.5	13.0	7.4
Alopecia	75.8	62.3	74.2
Asthenia	C1.0	52.5	66.2
Any Severe	61.8 12.8	52.5 24.6	66.3 14.9
Myalgia	12.0	24.0	14.3
Any	18.9	16.4	21.1
Severe	1.5	1.6	1.8
Arthralgia	9.2	6.6	8.2
Infusion Site Reactions	4.4	3.3	4.0

\*Normal Baseline LFTs: Transaminases ≤ 1.5 times ULN or alkaline phosphatase ≤ 2.5 times ULN or isolated elevations of transaminases or alkaline phosphatase up to 5 times ULN \*Elevated Baseline LFTs: SGOT and/or SGPT > 1.5 times ULN concurrent with alkaline phosphatase > 2.5 times ULN server with alkaline phosphatase > 2.5 times ULN concurrent with alkaline phosphatase concurrent with alkaline phosphatase concurrent with alkaline phosphatase concurrent with alkaline phosphatase up to 5 times ULN concurrent with alkaline phosphatase concurrent concurrent with alkaline phosphatase concurrent concurrent with alkaline phosphatase concurrent Hematologic: (see WARNINGS).
Reversible marrow suppression was the major dose-limiting toxicity of TAXOTERE. The median time to nadir was 7 days, while the median duration of severe neutropenia (~500 cells/mm³) was 7 days. Among 2045 patients with solid tumors and normal baseline LFTs, severe neutropenia occurred in 75.4% and lasted for more than 7 days in 2.9% of cycles. Febrile neutropenia (~500 cells/mm³) with fever > 38°C with IV antibiotics and/or hospitalization) occurred in 11% of patients with solid tumors, in 12.3% of patients with metastatic breast cancer, and in 9.8% of 92 breast cancer patients premedicated with 3-day corticosteroids. Severe infectious episodes occurred in 6.1% of patients with solid tumors, in 6.4% of patients with metastatic breast cancer, and in 5.4% of 92 breast cancer patients premedicated with 3-day corticosteroids.

Fluid Retention: (see BOXED WARNING, WARNINGS: Premedication Regimen, and PRECAU-

Cutaneous sections.

Cutaneous Severe skin toxicity is discussed in PRECAUTIONS. Reversible cutaneous reactions characterized by a rash including localized eruptions, mainly on the feet and/or hands, but also on the arms, face, or thorax, usually associated with pruritus, have been observed. Eruptions generally occurred within 1 week after TAXOTERE infusion, recovered before the next infusion, and were not disabling. Severe nail disorders were characterized by hypo- or hyperpigmentation, and occasionally by onycholysis (in 0.8% of patients with solid tumors) and pain.

Neurologic: (see PRECAUTIONS).

Gastrointestinal
Gastrointestinal reactions (nausea and/or vomiting and/or diarrhea) were generally mild to moderate. Severe reactions occurred in 3-5% of patients with solid tumors and to a similar extent among metastatic breast cancer patients. The incidence of severe reactions was 1% or less for the 92 breast cancer patients premedicated with 3-day corticosteroids. Severe stomatifis occurred in 5.5% of patients with solid tumors, in 7.4% of patients with metastatic breast cancer, and in 1.1% of the 92 breast cancer patients premedicated with 3-day corticosteroids.

Cardiovascular Hypotension occurred in 2.8% of patients with solid tumors; 1.2% required treatment. Clinically meaningful events such as heart failure, sinus tachycardia, atrial flutter, dysrhythmia, unstable angina, pulmonary edema, and hypertension occurred rarely, 8.1% (7/86) of metastatic breast cancer patients receiving TAXOTERE 100 mg/m² in a randomized trial and who had serial left ventricular ejection fractions assessed developed deterioration of LVEF by ≥ 10% associated with a drop below the institutional lower limit of normal. Infusion Site Reactions Infusion Site Reactions Infusion site reactions were generally mild and consisted of hyperpigmentation, inflammation, redness or dryness of the skin, phlebitis, extravasation, or swelling of the vein. Hepatic

Hepatic
In patients with normal LFTs at baseline, bilirubin values greater than the ULN occurred in 8.9% of patients. Increases in SGOT or SGPT > 1.5 times the ULN, or alkaline phosphatase > 2.5 times ULN, were observed in 18.9% and 7.3% of patients, respectively. While on TAXOTERE, increases in SGOT and/or SGPT > 1.5 times ULN concomitant with alkaline phosphatase > 2.5 times ULN occurred in 4.3% of patients with normal LFTs at baseline. (Whether these changes were related to the drug or underlying disease has not been established.) lematologic and Other Toxicity: Relation to dose and baseline liver chemistry abno Hematologic and other toxicity is increased at higher doses and in patients with elevated baseline liver function tests (LFTs). In the following tables, adverse drug reactions are compared for three populations: 730 patients with normal LFTs given TAXOTERE at 100 mg/m² in the randomized and single arm studies of metastatic breast cancer after failure of previous chemotherapy; 18 patients in these studies who had abnormal baseline LFTs (defined as SGOT and/or SGPT > 1.5 times ULN concurrent with alkaline phosphatase > 2.5 times ULN); and 174 patients in Japanese studies given TAXOTERE at 60 mg/m² who had normal LFTs.

Hematologic Adverse Events in Breast Cancer Patients
Previously Treated with Chemotherapy
Treated at TAXOTERE 100 mg/m² with Normal or Elevated Liver Function Tests or
60 mg/m² with Normal Liver Function Tests

		TAX0 100 n	TERE ng/m <sup>2</sup>	TAXOTERE 60 mg/m <sup>2</sup>
Adverse Eve	ent	Normal LFTs* n=730 %	Elevated LFTs** n=18 %	Normal LFTs* n=174 %
Neutropeni	a			
Any	<2000 cells/mm <sup>3</sup>	98.4	100	95.4
Grade 4	<500 cells/mm <sup>3</sup>	84.4	93.8	74.9
Thrombocy	topenia			
Any	<100,000 cells/mm <sup>3</sup>	10.8	44.4	14.4
Grade 4	<20,000 cells/mm <sup>3</sup>	0.6	16.7	1.1
Anemia	<11 g/dL	94.6	94.4	64.9
Infection**	*			
Any		22.5	38.9	1.1
Grade 3 ar		7.1	33.3	0
	ıtropenia****			
By Patient		11.8	33.3	0
By Course		2.4	8.6	0
<b>Septic Deat</b>	h	1.5	5.6	1.1
Non-Septic	Death	1.1	11.1	0
VLN or isola **Elevated phosphatas ***Incidence (n=62) amo	aseline LFTs: Transamin ated elevations of trans Baseline LFTs: SGOT ie >2.5 times ULN ce of infection requirin ing the 730 patients with nia, and 46 patients ha	saminases or alkaling and/or SGPT >1.5 g hospitalization an h normal LFTs at bas	e phosphatase up to times ULN concurr d/or intravenous an eline; 7 patients had	5 times ULN ent with alkaline ntibiotics was 8.5%

\*\*\*Febrile Neutropenia: For 100 mg/m², ANC grade 4 and fever > 38°C with IV antibiotics nd/or hospitalization; for 60 mg/m², ANC grade 3/4 and fever > 38.1°C Non-Hematologic Adverse Events in Breast Cancer Patients

TAXOTERE 100 mg/m²   TAXOTERE 60 mg/m²	Previously Treated with Chemotherapy  Treated at TAXOTERE 100 mg/m² with Normal or Elevated Liver Function Tests or  60 mg/m² with Normal Liver Function Tests						
Adverse Event		TAX	OTERE				
Reaction Regardless of Premedication	Adverse Event	LFTs* n=730	LFTs** n=18	LFTs* n=174			
Severe   1.2   0   0   0	Reaction Regardless of						
Regardless of Premedication							
Severe         7.9         16.7         0           Neurosensory Any Severe         56.8         50         19.5           Severe         5.8         0         0           Myalgia         22.7         33.3         3.4           Cutaneous Any Severe         44.8         61.1         30.5           Severe         4.8         16.7         0           Asthenia Severe         16.6         22.2         0           Diarrhea         0         0         0							
Any 56.8 50 19.5 Severe 5.8 0 0 0 Myalgia 22.7 33.3 3.4 Cutaneous 44.8 61.1 30.5 Severe 4.8 16.7 0 Severe 4.8 16.7 0 Asthenia Any 65.2 44.4 65.5 Severe 16.6 22.2 0 Diarrhea							
Severe         5.8         0         0           Myalgia         22.7         33.3         3.4           Cutaneous             Any             Any		E6.0	FO	10 F			
Cutaneous         44.8         61.1         30.5           Severe         4.8         16.7         0           Asthenia         Any         65.2         44.4         65.5           Severe         16.6         22.2         0           Diarrhea							
Any 44.8 61.1 30.5 Severe 4.8 16.7 0  Asthenia Any 65.2 44.4 65.5 Severe 16.6 22.2 0  Diarrhea	Myalgia	22.7	33.3	3.4			
Any 65.2 44.4 65.5 Severe 16.6 22.2 0  Diarrhea	Any						
	Any						
Severe 6.3 11.1	Any	42.2 6.3	27.8 11.1	NA			
Stomatitis         Any         53.3         66.7         19.0           Severe         7.8         38.9         0.6           *Normal Reseline LETs: Transaminases < 1.5 times LILN or alkaline phosphatase < 2.5 times	Any Severe	7.8	38.9	0.6			

\*Normal Baseline LFTs: Transaminases < 1.5 times ULN or alkaline phosphatase < 2.5 times ULN or isolated elevations of transaminases or alkaline phosphatase up to 5 times ULN \*\* Elevated Baseline Liver Function: SGOT and/or SGPT > 1.5 times ULN concurrent with alkaline phosphatase > 2.5 times ULN \*\*\*Fluid Retention includes (by COSTART): dema (peripheral, localized, generalized, lymphedema, pulmonary edema, and edema otherwise not specified) and effusion (pleural, pericardial, and ascites); no premedication given with the 60 mg/m² dose

In the three-arm monotherapy trial, TAX313, which compared TAXOTERE 60, 75 and 100 mg/m² in advanced breast cancer, the overall safety profile was consistent with the safety profile observed in previous TAXOTERE trials. Grade 3/4 or severe adverse events occurred in 49.0% of patients treated with TAXOTERE 60 mg/m² compared to 55.3% and 65.9% treated with 75 and 100 mg/m² respectively. Discontinuation due to adverse events was reported in 5.3% of patients treated with 60 mg/m² vs. 6.9% and 16.5% for patients treated at 75 and 100 mg/m² respectively. Deaths within 30 days of last treatment occurred in 4.0% of patients treated with 60 mg/m² compared to 5.3% and 1.6% for patients treated at 75 and 100 mg/m² respectively.

The following adverse events were associated with increasing docetaxel doses: fluid retention (26%, 38%, and 46% at 60, 75, and 100 mg/m² respectively), thrombocytopenia (7%, 11% and 12 % respectively), neutropenia (92%, 94%, and 97% respectively), febrile neutropenia (5%, 7%, and 14% respectively), treatment-related grade 3/ 4 infection (2%, 3%, and 7% respectively) and anemia (87%, 94%, and 97% respectively).

Combination Therapy with TAXOTERE in the Adjuvant Treatment of Breast Cancer The following table presents treatment emergent adverse events (TEAEs) observed in 744 paties who were treated with TAXOTERE 75 mg/m<sup>2</sup> every 3 weeks in combination with doxorubicin acyclophosphamide.

Clinically Important Treatment Emergent Adverse Events Regardless of Causal elationship in Patients Receiving TAXOTERE in Combination with Doxorubicin a Cyclophosphamide (TAX 316).

Cyclophosphamide (TAX 316).						
	TAXOTERE 75 mg/m²+ Doxorubicin 50 mg/m²+ Cyclophosphamide 500 mg/m² (TAC) n=744 %		Fluorouracil 500 mg/m²+ Doxorubicin 50 mg/m²+ Cyclophosphamide 500 mg/m² (FAC) n=736 %			
Adverse Event	Any	G 3/4	Any	G 3/4		
Anemia	91.5	4.3	71.7	1.6		
Neutropenia	71.4	65.5	82.0	49.3		
Fever in absence of infection	46.5	1.3	17.1	0.0		
Infection	39.4	3.9	36.3	2.2		
Thrombocytopenia	39.4	2.0	27.7	1.2		
Febrile neutropenia	24.7	N/A	2.5	N/A		
Neutropenic infection	12.1	N/A	6.3	N/A		
Hypersensitivity reactions	13.4	1.3	3.7	0.1		
Lymphedema	4.4	0.0	1.2	0.0		
Fluid Retention*	35.1	0.9	14.7	0.1		
Peripheral edema	26.9	0.4	7.3	0.0		
Weight gain	12.9	0.3	8.6	0.3		
Neuropathy sensory	25.5	0.0	10.2	0.0		
Neuro-cortical	5.1	0.5	6.4	0.7		
Neuropathy motor	3.8	0.1	2.2	0.0		
Neuro-cerebellar	2.4	0.1	2.0	0.0		
Syncope	1.6	0.5	1.2	0.3		
Alopecia	97.8	N/A	97.1	N/A		
Skin toxicity	26.5	0.8	17.7	0.4		
Nail disorders	18.5	0.4	14.4	0.1		
Nausea	80.5	5.1	88.0	9.5		
Stomatitis	69.4	7.1	52.9	2.0		
Vomiting	44.5	4.3	59.2	7.3		
Diarrhea	35.2	3.8	27.9	1.8		
Constipation	33.9	1.1	31.8	1.4		
Taste perversion	27.8	0.7	15.1	0.0		
Anorexia	21.6	2.2	17.7	1.2		
Abdominal Pain	10.9	0.7	5.3	0.0		
Amenorrhea	61.7	N/A	52.4	N/A		
Cough	13.7	0.0	9.8	0.1		
Cardiac dysrhythmias	7.9	0.3	6.0	0.3		
Vasodilatation	27.0	1.1	21.2	0.5		
Hypotension	2.6	0.0	1.1	0.1		
Phlebitis	1.2	0.0	0.8	0.0		
Asthenia	80.8	11.2	71.2	5.6		
Myalgia	26.7	0.8	9.9	0.0		
Arthralgia	19.4	0.5	9.0	0.3		
Lacrimation disorder	11.3	0.1	7.1	0.0		
Conjunctivitis	5.1	0.3	6.9	0.1		

\* COSTART term and grading system for events related to treatment. Of the 744 patients treated with TAC, 36.3% experienced severe TEAEs compared to 26.6 % of the 736 patients treated with FAC. Dose reductions due to hematologic toxicity occurred in 1% of cycles in the TAC

being the most common reasons for withdrawal among TAC-treated patients. Two patients died in each arm within 30 days of their last study treatment; 1 death per arm was attributed to study drugs. Fever and Infection
Fever in the absence of infection was seen in 46.5% of TAC-treated patients and in 17.1% of FAC-treated

arm versus 0.1% of cycles in the FAC arm. Six percent of patients treated with TAC discontinued treatmendue to adverse events, compared to 1.1% treated with FAC; fever in the absence of infection and allergy

TAC-treated patients required treatment discontinuation; no deaths due to these events occurred. More cardiovascular events were reported in the TAC arm vs. the FAC arm: dvsrhvthmias, all grades (7.9%)

vs. 6.0%), hypotension, all grades (2.6% vs. 1.1%) and CHF (1.6% vs. 0.5%). One patient in each arm died Acute Myeloid Leukemia Treatment-related acute myeloid leukemia (AML) is known to occur in patients treated with anthracy-clines and/or cyclophosphamide, including use in adjuvant therapy for breast cancer. AML occurs at a higher frequency when these agents are given in combination with radiation therapy. AML occurred in the adjuvant breast cancer trial (TAX316). The cumulative risk of developing treatment-related AML at 5 years in TAX316 was 0.4% for TAC-treated patients and 0.1% for FAC-treated patients. This risk of AML is comparable to the risk observed for other anthracyclines/cyclophosphamide containing adjuvant breast chemotherapy regimes.

oreast chemotherapy regimens.
Woonotherapy with TAXOTERE for Unresectable, Locally Advanced or Metastatic NSCLC Previously Treated with Platinum-Based Chemotherapy TAXOTERE 75 mg/m<sup>2</sup>: Treatment emergent adverse drug reactions are shown below. Included in this table are safety data for a total of 176 patients with non-small cell lung carcinoma and a history of prior treatment with platinum-based chemotherapy who were treated in two randomized, controlled trials. These reactions were described using NCI Common Toxicity Criteria regardless of relationship to study treatment, except for the hema

Treatment Emergent Adverse Events Regardless of Relationship to Treatment in Patients Receiving TAXOTERE as Monotherapy for Non-Small Cell Lung Cancer Previously

	TAXOTERE	Sed Chemotherapy*  Best Supportive	Vinorelbine/
	75 mg/m <sup>2</sup>	Care	Ifosfamide
	n=176	n=49	n=119
Adverse Event	%	%	%
Neutropenia	04.4	442	02.2
Any Grade 3/4	84.1 65.3	14.3 12.2	83.2 57.1
Leukopenia	03.3	12,2	37.1
Any	83.5	6.1	89.1
Grade 3/4	49.4	0	42.9
Thrombocytopenia	0.0		7.6
Any Grade 3/4	8.0 2.8	0	7.6 1.7
Anemia	2.0	0	1.7
Any	91.0	55.1	90.8
Grade 3/4	9.1	12.2	14.3
Febrile			
Neutropenia**	6.3	NA†	0.8
Infection Any	33.5	28.6	30.3
Grade 3/4	10.2	6.1	9.2
Treatment Related Mortality	2.8	NA†	3.4
Hypersensitivity Reactions			
Any	5.7	0	0.8
Grade 3/4	2.8	0	0
Fluid Retention	33.5	ND††	22.7
Any Severe	2.8	NUIT	3.4
Neurosensory	2.0		5.1
Any	23.3	14.3	28.6
Grade 3/4	1.7	6.1	5.0
Neuromotor			
Any	15.9	8.2	10.1
Grade 3/4	4.5	6.1	3.4
Skin Any	19.9	6.1	16.8
Grade 3/4	0.6	2.0	0.8
Gastrointestinal	0.0	2.0	0.0
Nausea			
Any	33.5	30.6	31.1
Grade 3/4	5.1	4.1	7.6
Vomiting Any	21.6	26.5	21.8
Grade 3/4	2.8	2.0	5.9
Diarrhea		2.0	3.3
Any	22.7	6.1	11.8
Grade 3/4	2.8	0	4.2
Alopecia	56.3	34.7	49.6
Asthenia	52.0	57.1	52.0
Any Severe***	52.8 18.2	57.1 38.8	53.8 22.7
Stomatitis	10.2	30.0	44.1
Any	26.1	6.1	7.6
Grade 3/4	1.7	0	0.8
Pulmonary			
Any	40.9	49.0	45.4
Grade 3/4	21.0	28.6	18.5
Nail Disorder Any	11.4	0	1.7
Severe***	1.1	0	0
Myalgia		Ť	
Anv	6.3	0	2.5
Severe***	0	0	0
Arthralgia	-		
Any Severe***	3.4	2.0	1.7 0.8
	0	0	υ.δ
Taste Perversion Any	5.7	0	0
Severe***	0.6	0	0
	0.0		

\*Normal Baseline LFTs: Transaminases ≤ 1.5 times ULN or alkaline phosphatase ≤ 2.5 times ULN or isolated elevations of transaminases or alkaline phosphatase up to 5 times ULN \*\*Febrile Neutropenia: ANC grade 4 with fever > 38°C with IV antibiotics and/or hospitalization \*\*\*COSTART term and grading system
† Not Applicable; †† Not Done

ation Therapy with TAXOTERE in Chemotherapy-Naïve Advanced Unresectable or The table below presents safety data from two arms of an open label, randomized controlled trial (TAX326) that enrolled patients with unresectable stage IIIB or IV non-small cell lung cancer and no history of prior chemotherapy. Adverse reactions were described using the NCI Common Toxicity Criteria except where otherwise noted.

Neutropenia	TAXOTERE 75 mg/m² + Cisplatin 75 mg/m² n=406	Vinorelbine 25 mg/m² + Cisplatin 100 mg/m² n=396 %
Grade 3/4         74         78           Febrile Neutropenia         5         5           Thrombocytopenia         3         4           Any         15         15           Grade 3/4         3         4           Annemia         3         4           Any         89         94           Grade 3/4         7         25           Infection         3         8           Any         35         37           Grade 3/4         8         8           Fever in absence of infection         3         29           Any         3         29           Grade 3/4         <1         1           Hypersensitivity Reaction*         3         29           Any         12         4         4           Any         12         4         4           Any         12         4         4           All severe or life-threatening events         2         2         2           Petripheral edema         2         2         2         2         2           Any         3         4         18         18         18         18         18		
Series   S		
Thrombocytopenia	1.1	
Any	5	5
Grade 3/4         3         4           Anemia         89         94           Grade 3/4         7         25           Infection         35         37           Any         35         37           Grade 3/4         8         8           Fever in absence of infection         3         29           Grade 3/4         <1	15	15
Any Grade 3/4 7 25  Infection Any 35 37  Grade 3/4 8 8 8 8  Rever in absence of infection Any 33 29  Grade 3/4		
Grade 3/4		
Infection		
Any Grade 3/4 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	/	23
Fever in absence of infection Any 33 29 Grade 3/4 < 1 1 Hypersensitivity Reaction* Any 12 4 Grade 3/4 3		
Any Grade 3/4	8	8
State   Stat	22	20
Hypersensitivity Reaction*		
Any Grade 3/4		<u> </u>
Fluid Retention**  Any 54 42 All severe or life-threatening events 2 2 2 Pleural effusion Any 23 22 All severe or life-threatening events 2 2 2 Penipheral edema Any 34 18 All severe or life-threatening events 2 2 2 Penipheral edema Any 34 18 All severe or life-threatening events 34 18 All severe or life-threatening events 34 34 34 34 34 34 34 34 34 34 34 34 34		
Any	3	<1
All severe or life-threatening events Pleural effusion Any All severe or life-threatening events 2 2 2 2 2 2 3 3 22 3 3 22 3 3 3 3 4 3 4 3 3 4 3 3 4 3 3 4 3 3 4 3 3 4 3 3 4 3 3 4 3 3 4 3 3 4 3 3 4 3 4 3 3 4 3 4 3 3 4 3 4 3 3 4 3 4 3 3 4 3 4 3 3 4	54	42
Any All severe or life-threatening events 2 2 2 2 Peripheral edema Any 34 18 18		
All severe or life-threatening events Any All severe or life-threatening events  Neurosensory Any Any Afrade 3/4 Any All severe or life-threatening events  Any Any Any Any Any Any Any Any Any An	22	
Peripheral edema Any All severe or life-threatening events Weight gain Any All severe or life-threatening events  Any All severe or life-threatening events  Self and any A		
Any All severe or life-threatening events	2	2
Weight gain		
Any	<1	<1
All severe or life-threatening events    Neurosensory   Any	15	9
Any		
Grade 3/4  Neuromotor  Any Grade 3/4  Any Grade 3/4  Any Any Grade 3/4  Any Any Any Any Ary Ary Ary Ary Any Ary Ary Ary Ary Ary Ary Ary Ary Ary Ar		
Neuromotor		
Any 6	4	7
Skin Any Any Any Any Any Any Any Any Any An		17
Any	3	6
Grade 3/4 <1 1  Nausea Any 72 76 Grade 3/4 10 17  Vomiting Any 55 61 Grade 3/4 8 16  Diarrhea Any 47 25 Grade 3/4 7 3  Anorexia** Any 42 40 All severe or life-threatening events 5 5  Stomatitis Any 24 21 Grade 3/4 2 1  Alopecia Any 24 21 Grade 3/4 2 1  Alopecia Any 75 42 Alopecia Any 75 44 All severe or life-threatening events 12 14  Nail Disorder** Any 14 <1 All severe events <1 0  Myalgia** Any 18 12	16	14
Nausea     72     76       Grade 3/4     10     17       Vomiting     55     61       Any     55     61       Grade 3/4     8     16       Diarrhea     7     3       Any     47     25       Grade 3/4     7     3       Anorexia**     40     41       Any est ever or life-threatening events     5     5       Stomatitis     2     40       Any     24     21       Grade 3/4     2     1       All severe and any     75     42       Grade 3     <1		1.1
Grade 3/4     10     17       Vomiting     55     61       Any     55     61       Diarrhea     7     3       Any     47     25       Grade 3/4     7     3       Anorexia**     42     40       All severe or life-threatening events     5     5       Stomatitis     24     21       Grade 3/4     2     1       Alopecia     75     42       Grade 3     <1		
Vomiting         4ny         55         61         62         61         61         62         62         61         62         62         62         63         64         62         64         63         64		
Any 55 61 Grade 3/4 8 16  Diarrhea Any 47 25 Grade 3/4 7 3  Anorexia** Any 42 40 All severe or life-threatening events 5 5  Stomatitis Any 24 21 Grade 3/4 2 1  Alopecia Any 75 42 Grade 3 <1 0  Asthenia** Any 74 75 All severe or life-threatening events 12 14  Nail Disorder** Any 75 All severe or life-threatening events 12 14  Mail Store or life-threatening events 12 14  May 75 All severe or life-threatening events 12 14  May 14 < 1 All severe events <1 0  Myalgia** Any 18 12	10	17
Grade 3/4  B 16  Diarrhea  Any 47 25  Grade 3/4 7 3  Anorexia**  Any 42 40  All severe or life-threatening events 5  Stomatitis  Any 24 21  Grade 3/4 2 1  All pecia  Any 75 42  Grade 3 4 0  Asthenia**  Any 75 42  Asthenia**  Any 74 75  All severe or life-threatening events 12  Any 75 42  Asthenia**  Any 74 75  All severe or life-threatening events 12  Any 74 75  All severe or life-threatening events 12  Any 74 75  All severe or life-threatening events 12  Any 74 75  Any 75  Any 74 75  Any 74 75  Any 75  Any 74 75  Any 75  Any 74 75  Any 74 75  Any 75  Any 75  Any 74 75  Any 75	55	61
Any 47 25 Grade 3/4 7 3  Anorexia** Any 42 40 All severe or life-threatening events 5 5  Stomatitis Any 24 21 Grade 3/4 2 1  Alopecia Any 75 42 Grade 3 < 1 0  Asthenia** Any 74 75 All severe or life-threatening events 12 14  Nail Disorder** Any 74 < 75 All severe events < 1 0  Myalgia** Any 18 12		
Grade 3/4     7     3       Anorexia**     42     40       Any     42     40       All severe or life-threatening events     5     5       Stomatitis     3     24     21       Grade 3/4     2     1     1       Alopecia     42     42     6       Any     75     42     6       Grade 3     <1		
Anorexia**  Any 42 40 All severe or life-threatening events 5 5  Stomatitis  Any 24 21 Grade 3/4 2 1  Alopecia  Any 75 42 Grade 3 < 1 0  Asthenia**  Any 74 75 All severe or life-threatening events 12 14  Nail Disorder**  Any 14 < 1  All severe events < 1 0  Myalgia**  Any 18 12		
Any All severe or life-threatening events 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	/	<u> </u>
Stomatitis	42	40
Any 24 21 Grade 3/4 2 1  Alopecia Any 75 42 Grade 3 < 1 0  Asthenia** Any 74 75 All severe or life-threatening events 12 14  Nail Disorder** Any 14 < 1 All severe events < 1 0  Myalgia** Any 18 12	5	5
Grade 3/4 2 1  Alopecia Any 75 42 Grade 3 <1 0  Asthenia** Any 74 75 All severe or life-threatening events 12 14  Nail Disorder** Any 14 <1 All severe events <1 0  Myalgia** Any 18 12	24	21
Alopecia     75     42       Any     75     42       Grade 3     <1		
Ány     75     42       Grade 3     <1	_	
Asthenia**  Any 74 75 All severe or life-threatening events 12 14  Nail Disorder**  Any 14 < 1  All severe events <1 0  Myalgia**  Any 18 12		
Any 74 75 All severe or life-threatening events 12 14  Nail Disorder**  Any 14 < 1 All severe events <1 0  Myalgia**  Any 18 12	<1	0
All severe or life-threatening events 12 14  Nail Disorder** Any 14 < 1  All severe events <1 0  Myalgia** Any 18 12	74	75
Nail Disorder**       Any     14     <1		
All severe events <1 0  Wyalgia** Any 18 12		
<b>Myalgia**</b> Any 18 12		
Any 18 12	<1	0
	18	12
All severe events <1 <1	<1	<1
* Replaces NCI term "Allergy"		75 mg/m² + Cisplatin 75 mg/m² n=406 %  91 74 5  15 3 89 7 35 88 33 <1 12 33 <41 15 <47 4 4 19 3 16 <41 17 47 4 19 3 16 <41 72 10 55 8 47 7 42 5 24 27 75 <41 74 12 14 <41 18

docetaxel+cisplatin arm and 37 patients (9.3%) in the vinorelibine+cisplatin arm. Deaths within 30 days of last study treatment attributed to study drug occurred in 9 patients (2.2%) in the docetaxel+cisplatin arm and 8 patients (2.0%) in the vinorelibine+cisplatin arm. not demonstrate a superior survival associated with TAXOTERE, see CLINICAL STUDIES section or definition and a higher incidence of thrombocytopenia, diarrhea, fluid retention, hypersensitivi actions, skin toxicity, alopecia and nail changes on the TAXOTERE+carboplatin arm, while a high cidence of anemia, neurosensory toxicity, nausea, vomiting, anorexia and asthenia was observed or a strength of the control o

bination Therapy with TAXOTERE in Patients with Prostate Cancer The following data are based on the experience of 332 patients, who were treated with TAXOTERE 75 mg/m² every 3 weeks in combination with prednisone 5 mg orally twice daily. (docetaxel) Injection Concentrate

TAXOTERE 75 mg/m² every 3 weeks + prednisone 5 mg twice daily n=332		Mitoxantrone 12 mg/m² every 3 weeks + prednisone 5 mg twice daily n=335	
G 3/4	Any	G 3/4	
4.9	57.8	1.8	
32.0	48.2	21.7	
0.6	7.8	1.2	
N/A	1.8	N/A	
5.7	20.3	4.2	
0.3	1.8	0.0	
0.6	0.6	0.0	
0.6	4.5	0.3	
0.3	3.0	0.0	
0.3	1.5	0.0	
1.8	7.2	0.3	
1.5	3.0	0.9	
0.3	3.3	0.6	
N/A	12.8	N/A	
0.0	7.5	0.0	
2.7	35.5	1.5	
2.1	9.6	1.2	
0.9	8.4	0.0	
0.0	6.6	0.0	
1.5	14.0	1.5	
1.2	14.3	0.3	
0.0	7.8	0.0	
2.7	8.7	0.9	
0.3	22.1	1.2	
4.5	34.6	5.1	
		0.9	
		0.9	
		1.2	
	0.3 0.6 0.6	0.6 1.5	

Post-marketing Experiences lance. Because they are reported from a population of unknown size, pre

Body as a whole: diffuse pain, chest pain, radiation recall pher Cardiovas unlar: atrial fibrillation, deep vien thrombosis, EEG abnormalities, thrombophlebitis, pulmonary embolism, syncope, tachycardia, myocardial infarction Cutaneous: rare cases of bullous eruption such as erythema multiforme or Stevens-Johnson Itinle factors may have contributed to the development of these effects testinal: abdominal pain, anorexia, constipation, duodenal ulcer, esophagitis, gastroin hemorrhage, gastrointestinal perforation, ischemic colitis, colitis, intestinal obstruction utropenic enterocolitis and dehydration as a consequence to gastrointestinal events have

curriogic: confusion, rare cases of seizures or transient loss of consciousness have been observed, memory confusion, rare cases of seizures or transient loss of consciousness have been observed, the drug, confusion of the confusion of the drug. tearing which may be attributable to lacrimal duct obstruction has been reported. Rar cases of transient visual disturbances (flashes, flashing lights, scotomata) typically occurring during drug infusion and in association with hypersensitivity reactions have been reported. These were Respiratory: dyspnea, acute pulmonary edema, acute respiratory distress syndrome, interstitial pneumonia. Pulmonary fibrosis has been rarely reported.

ere is no known antidote for TAXOTERE overdosage. In case of overdosage, the patient should be overdosage include: bone marrow suppression, peripheral neurotoxicity, and mucositis. Patients tould receive therapeutic G-CSF as soon as possible after discovery of overdose. Other appropriate two reports of overdose, one patient received 150 mg/m<sup>2</sup> and the other received 200 mg/m<sup>2</sup> as -hour infusions. Both patients experienced severe ne thality was observed following single IV doses that were ≥154 mg/kg (about 4.5 times the ided human dose on a mg/m² basis); neurotoxicity associated with paralysis, non-extend limbs, and myelin degeneration was observed in mice at 48 mg/kg (about 1.5 times the ided human dose on a mg/m² basis). In male and female rats, lethality was observed at a nmended human dose on a mg/m<sup>2</sup> basis) and was asso ormal mitosis and necrosis of multiple organs.

### DOSAGE AND ADMINISTRATION

cisplatin 75 mg/m<sup>2</sup> over 30-60 minutes every 3 weeks.

nmended dose of TAXOTERE is 60-100 mg/m<sup>2</sup> administered intravenously over 1 hour even In the adjuvant treatment of operable node-positive breast cancer, the recommended TAXOTERE dose is 75 mg/m² administered 1-hour after doxorubicin 50 mg/m² and cyclophosphamide 500 mg/m² every 3 weeks for 6 courses. Prophylactic G-CSF may be used to mitigate the risk of ies (see also **Dosage Adjustments**). Non-Small Cell Lung Cancer

monotherapy, and the recommended dose is 75 mg/m<sup>2</sup> administered intravenously over 1 hour every 3 weeks. A dose of 100 mg/m<sup>2</sup> in patients previously treated with chemotherapy was associated with related mortality in randomized, controlled rials (see BOXED WARNING, WARNINGS and CLINICAL STUDIES sections). nistered intravenously over 1 hour immediately followed

Prostate cancer very 3 weeks as a 1 hour infusion. Prednisone 5 mg orally twice daily is adm Il patients should be premedicated with oral corticosteroids (see below for prostate cancer) such as methasone 16 mg per day (e.g., 8 mg BID) for 3 days starting 1 day prior to TAXOTERE adm vity reactions (see BOXED WARNING, WARNINGS, and PRECAUTIONS sect one 8 mg, at 12 hours, 3 hours and 1 hou hefore the TAXOTERE infusion (see WARNINGS, and PRECAUTIONS s

ents who are dosed initially at 100 mg/m<sup>2</sup> and who experience either febrile neutropeni Patients who are dosed initially at 100 mg/m² and who experience either febrile neutropenia, neutrophis <500 cells/mm³ for more than 1 week, or severe or cumulative cutaneous reactions during TAXOTERE therapy should have the dosage adjusted from 100 mg/m² to 75 mg/m². If the patient continues to experience these reactions, the dosage should either be decreased from 75 mg/m² to 55 mg/m² or the treatment should be discontinued. Conversely, patients who are dosed initially at 60 mg/m² and who do not experience febrile neutropenia, neutrophils <500 cells/mm³ for more than 1 week, severe or cumulative cutaneous reactions, or severe peripheral neuropathy duri TAXOTERE therapy may tolerate higher doses. Patients who develop ≥ grade 3 peripheral neuropat

ation Therapy with TAXOTERE in the Adjuvant Treatment of Breast Cancer rophil count is ≥ 1,500 cells/mm³. Patients who experience febrile neutropenia should recei -CSF in all subsequent cycles. Patients who continue to experience this reaction should remain or -CSF and have their TAXOTERE dose reduced to 60 mg/m<sup>2</sup>. Patients who experience severe or cumula tive cutaneous reactions or moderate neurosensory signs and/or symptoms during TAXOTERE therapy should have their dosage of TAXOTERE reduced from 75 to 60 mg/m². If the patient continues to experi Non-Small Cell Lung Cancer

Monotherapy with IAXOLEKE for MSLEL Tredument Pyter frame by First Tradition Postal Chambers Patients who are dosed initially at 75 mg/m² and who experience either febrile neutropenia, neutrophils <500 cells/mm³ for more than one week, severe or cumulative cutaneous reactions, or other grade 3/4 non-hematological toxicities during TAXOTERE treatment should have treatment withheld until resolution of the toxicity and then resumed at 55 mg/m². Patients who develop ≥ grade 3 peripheral neuropathy should have TAXOTERE treatment disco Combination Therapy with TAXOTERE for Chemotherapy-Naïve NSCLC
For patients who are dosed initially at TAXOTERE 75 mg/m<sup>2</sup> in combination with cisplatin, and ose nadir of platelet count during the previous course of therap tients who experience febrile neutropenia, and in patients with seri

ties, the TAXOTERE dosage in subsequent cycles should be reduced to 65 mg/m<sup>2</sup>. In patirequire a further dose reduction, a dose of 50 mg/m<sup>2</sup> is recommended. For cisplatin dosage tion Therapy with TAXOTERE for Hormone-Refractory Metastatic Prostate Cancer raxot Ext. Should be administed when the neutropin count is 2 1,500 censymbr. Fallens will experience either febrile neutropenia, neutrophils < 500 cells/mm³ for more than one week, sever or cumulative cutaneous reactions or moderate neurosensory signs and/or symptoms durin TAXOTERE therapy should have the dosage of TAXOTERE reduced from 75 to 60 mg/m². If the patier

rment: Patients with bilirubin > ULN should generally not receive TAXOTERE. Also patients with SGOT and/or SGPT > 1.5 x ULN concomitant with alkaline phosphatase > 2.5 x ULNshould generally not receive TAXOTERE.

Children: The safety and effectiveness of docetaxel in pediatric patients below the age of 16 years Elderly: See **Precautions, Geriatric Use.** In general, dose selection for an elderly patient should be cautious, reflecting the greater frequency of decreased hepa concomitant disease or other drug therapy in elderly patients.

es to experience these reactions at 60 mg/m<sup>2</sup>, the treatment should be discont

## PREPARATION AND ADMINISTRATION

Administration Precautions
TAXOTERE is a cytotoxic anticancer drug and, as with other potentially toxic compounds, caution should be exercised when handling and preparing TAXOTERE solutions. The use of gloves is recommended. Please refer to Handling and Disposal section.
If TAXOTERE Injection Concentrate, initial diluted solution, or final dilution for infusion should come into contact with the skin, immediately and thoroughly wash with soap and water. If TAXOTERE Injec-tion Concentrate, initial diluted solution, or final dilution for infusion should come into contact with

mucosa, immediately and thoroughly wash with water.

Contact of the TAXOTERE concentrate with plasticized PVC equipment or devices used to prepare olutions for infusion is not recommended. In order to minimize patient exposure to the plasticizer DEHP (di-2-ethylhexyl phthalate), which may be leached from PVC infusion bags or sets, the final TAXOTERE dilution for infusion should be stored in bottles (glass, polypropylene) or plastic bags (polypropylene, polyolefin) and administered through polyethylene-lined administration sets. TAXOTERE Injection Concentrate requires two dilutions prior to administration. Please follow the preparation instructions provided below. Note: Both the TAXOTERE Injection Concentrate and the dilutent vials contain an overfill to compensate for liquid loss during preparation. This overfill ensures that after dilution with the <a href="entity">entity</a> contents of the accompanying diluent, there is an initial diluted solution Containing 10 mg/mL docetaxel.

The table below provides the fill range of the diluent, the approximate extractable volume of diluen when the entire contents of the diluent vial are withdrawn, and the concentration of the initia rawn, and the concentration of the initial diluted solution for TAXOTERE 20 mg and TAXOTERE 80 mg.

duct	Diluent 13% (w/w) ethanol in water for injection Fill Range (mL)	Approximate extractable volume of diluent when entire contents are withdrawn (mL)	Concentration of the initial diluted solution (mg/mL docetaxel)		
otere® ng/0.5 mL	1.88 – 2.08 mL	1.8 mL	10 mg/mL		
otere® ng/2 mL	6.96 - 7.70 mL	7.1 mL	10 mg/mL		
aration and Administration					

A. Initial Diluted Solution

1. TAXOTERE vials should be stored between 2 and 25°C (36 and 77°F). If the vials are stored under refrigeration, allow the appropriate number of vials of TAXOTERE Injection Concentrate and diluent (13% ethanol in water for injection) vials to stand at room temperature for approximately

3 minutes.
2. Aseptically withdraw the entire contents of the appropriate diluent vial (approximately 1.8 mL for TAXOTERE 20 mg and approximately 7.1 mL for TAXOTERE 80 mg) into a syringe by partially inverting the vial, and transfer it to the appropriate vial of TAXOTERE Injection Concentrate. If the procedure is followed as described, an initial diluted solution of 10mg docetaxel/mL will result. 3. Mix the initial diluted solution by repeated inversions for at least 45 seconds to assure full mixture of the concentrate and diluent. Do not shake.

4. The initial diluted TAXOTERE solution (10 mg docetaxel/mL) should be clear; however, there may be some foam on top of the solution due to the polysorbate 80. Allow the solution to stand for a few minutes to allow any foam to dissipate. It is not required that all foam dissipate prior to continuing the preservation process.

continuing the preparation process. The initial diluted solution may be used immediately or stored either in the refrigerator or at

The initial diluted solution may be used immediately or stored either in the refrigerator or at room temperature for a maximum of 8 hours.

8. Final Dilution for Infusion

1. Aseptically withdraw the required amount of initial diluted TAXOTERE solution (10 mg docetaxel/mL) with a calibrated syringe and inject into a 250 mL infusion bag or bottle of either 0.9% Sodium Chloride solution or 5% Dextrose solution to produce a final concentration of 0.3 to 0.74 mg/mL If a dose greater than 200 mg of TAXOTERE is required, use a larger volume of the infusion vehicle so that a concentration of 0.74 mg/mL TAXOTERE is not exceeded.

2. Thoroughly mix the infusion by manual rotation.

3. As with all parenteral products, TAXOTERE should be inspected visually for particulate matter or discoloration prior to administration whenever the solution and container permit. If the TAXOTERE initial diluted solution or final dilution for infusion is not clear or appears to have precipitation, these should be discarded.

The final TAXOTERE dilution for infusion should be administered intravenously as a 1-hour infusion under ambient room temperature and lighting conditions.

Stability

Stability
TAXOTERE infusion solution, if stored between 2 and 25°C (36 and 77°F) is stable for 4 hours. Fully prepared TAXOTERE infusion solution (in either 0.9% Sodium Chloride solution or 5% Dextrose solution) should be used within 4 hours (including the 1 hour i.v. administration).

TAXOTERE Injection Concentrate is supplied in a single-dose vial as a sterile, pyrogen-free, non-aqueous, viscous solution with an accompanying sterile, non-pyrogenic, Diluent (13% ethanol in water for injection) vial. The following strengths are available:

TAXOTERE 80 mg/2 ML (NDC 0075-8001-80)

TAXOTERE (doctaxel) Injection Concentrate 80 mg/2 mt. 80 mg docetaxel in 2 mL polysorbate 80 and Diluent for TAXOTERE 80 mg (13% (w/w) ethanol in water for injection). Both items are in a blister

TAXOTERE 20 mg/0.5 ML (NDC 0075-8001-20)

TAXOTERE (docetaxel) Injection Concentrate 20 mg/0.5 mL: 20 mg docetaxel in 0.5 mL polysorbate 80 and diluent for TAXOTERE 20 mg (13% (w/w) ethanol in water for injection). Both items are in a blister

pack in one carion.

Storage

Store between 2 and 25°C (36 and 77°F). Retain in the original package to protect from bright light.
Freezing does not adversely affect the product.

Handling and Disposal

Procedures for proper handling and disposal of anticancer drugs should be considered. Several guidelines on this subject have been published¹-7. There is no general agreement that all of the procedures recommended in the guidelines are necessary or appropriate.

REFERENCES

1. OSHA Work-Practice Guidelines for Controlling Occupational Exposure to Hazardous Drugs. Am J Health-Syst Pharm. 1996; 53: 1669-1685.

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ventis Pharmaceuticals Inc ridgewater, NJ 08807 USA

(docetaxel) Injection Concentrate Dexamethasone tablets may protect patients from significant fluid retention. It is important that you take this medicine on

Taxotere treatment. Gastrointestinal - Diarrhea has been associated with TAXOTERE use and can be severe in some patients. Nausea and/or vomiting are common in patients receiving TAXOTERE. Severe inflammation of the bowel can also occur in some patients and may be life threatening.

schedule. If you have not taken dexamethasone on schedule,

you must tell your doctor or nurse before receiving your next

**Hair Loss** – Loss of hair occurs in most patients taking Taxotere (including the hair on your head, underarm hair, pubic hair, eyebrows, and eyelashes). Hair loss will begin after the first few treatments and varies from patient to patient. Once you have completed all your treatments, hair generally grows back.

Your doctor or nurse can refer you to a store that carries wigs, hairpieces, and turbans for patients with cancer.

**Fatigue** – A number of patients (about 10%) receiving Taxotere feel very tired following their treatments. If you feel tired or weak, allow yourself extra rest before your next treatment. If it is bothersome or lasts for longer than 1 week, inform your doctor or nurse.

**Muscle Pain** – This happens about 20% of the time, but is rarely severe. You may feel pain in your muscles or joints. Tell your doctor or nurse if this happens. They may suggest ways to make you more comfortable.

**Rash** – This side effect occurs commonly but is severe in about 5%. You may develop a rash that looks like a blotchy, hive-like reaction. This usually occurs on the hands and feet but may also appear on the arms, face, or body. Generally, it will appear between treatments and will go away before the next treatment. Inform your doctor or nurse if you experience a rash. They can help you avoid discomfort.

**Odd Sensations** – About half of patients getting Taxotere will feel numbness, tingling, or burning sensations in their hands and feet. If you do experience this, tell your doctor or nurse. Generally, these go away within a few weeks or months after your treatments are completed. About 14% of patients may also develop weakness in their hands and feet.

**Nail Changes** – Color changes to your fingernails or toenails may occur while taking Taxotere. In extreme, but rare, cases nails may fall off. After you have finished Taxotere treatments, your nails will generally grow back.

**Eye Changes** – Excessive tearing, which can be related to conjunctivitis or blockage of the tear ducts, may occur.

If you are interested in learning more about this drug, ask your doctor for a copy of the package insert.

**Aventis Pharmaceuticals Inc.** 

Bridgewater, NJ 08807 USA www.aventis-us.com

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**Every three-week injection of TAXOTERE for** breast and non-small cell lung cancers Take dexamethasone tablets, 8 mg twice daily. **Dexamethasone dosing: Day 1** Date: \_\_\_\_\_Time: \_\_\_\_AM\_\_\_\_PM **Day 2** Date: \_\_\_\_\_ \_\_Time:\_\_\_\_ (Taxotere Treatment Day) **Day 3** Date: \_\_\_\_\_\_Time: \_\_\_\_\_AM\_\_\_\_\_PM

prostate cancer Take dexametha	k injection of TAXOTERE for sone 8 mg, at 12 hours, our before TAXOTERE infusion.
Dexamethasone	dosing:
Date:	Time:
Date:	Time:
(Taxotere Treatmer	nt Day)
	Time: